



CONFORMITY TO TYPE BASED ON INTERNAL PRODUCTION CONTROL PLUS SUPERVISED PRODUCT CHECKS AT RANDOM INTERVALS (MODULE C2)

MODÜL C2 - ÜRETİMİN DÂHİLÎ KONTROLÜ VE ÜRÜNÜN RASTGELE ARALIKLARLA DENETİMLİ MUAYENESİNE DAYALI TİPE UYGUNLUK

: 27092537

: 10.10.2024-10.10.2025

Belge No / Certificate No

Belgelendirme Tarihi - Bir Sonraki Belge Tarihi /

Certification Date / Certificate Validity Date

Belge Geçerlilik Tarihi / Document Validity Period : 1 yıl / 1 year

Firma Unvanı ve Adresi /

Company Name and Address : JEDX MEDCARE

Köysikuja 1, 01640 Vantaa, FINLAND

Marka / Model / Brand / Model : JEDX 1625 W HL

Direktifi / Directive : 2016/425 REGULATION

Modülü/Kategori / Module / Category : C2 MODÜLÜ/ KATEGORİ III

MODULE C2 / CATEGORY III

Teknik Değerlendirme Rapor No/

Technical Evaluation Report No : 27092537

Ürün Tipi / Product Type:

- EN 149:2001+ A1:2009 Solunumla ilgili koruyucu cihazlar - Parçacıklara karşı koruma amaçlı filtreli yarım maskeler/ *Respiratory protective devices - Filtering half masks to protect against particles*

Ürünün Malzeme Bilgisi / *Product Material Information*: JEDX 1625 W HL model ürünleri kumaş, elastik kayış, burun klipsi ve filtre katmanı kullanılarak imal edilmiştir./ JEDX 1625 W HL model products are manufactured using fabric, elastic strap, nose clip, filter layer.

Karar Verici / Approver Şirket Müdürü / General Manager

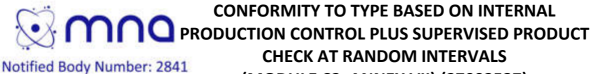


MNA Laboratuvarları San. Tic.Ltd .Şti Adres: Küçükbakkalköy Mahallesi Yenidoğan Cad.No:21 Ataşehir/ İstanbul Tel: 0216 574 07 08 Faks: 0216 575 13 31 www.mnalab.com

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CHECK AT RANDOM INTERVALS (MODULE C2, ANNEX VII) (27092537)

Report No : 27092537

Report Date : 10.10.2024

Application No : 27092537

1. COMPANY INFORMATION:

JEDX MEDCARE

Köysikuja 1, 01640 Vantaa, FINLAND

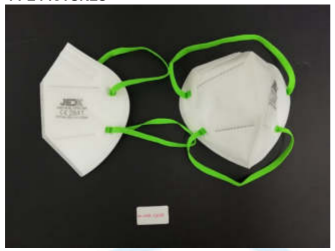
2. PPE INFORMATION:

Disposable and non-sterile half mask made of particulate protection fitler material.

3. PPE TYPE IDENTIFICATION

EN 149:2001+A1:2009 Respiratory protective devices - Filtering half masks to protect against particles -Requirements, testing, marking

4. PPE PICTURES



JEDX 1625 W HL

5. PPE DIMENSIONS:

JEDX 1625 W HL model has been found to be produced using standard sizes.

6. PPE PRODUCT MATERIAL INFORMATION:

The mask is made of elastic strap, nonwoven fabric on the outer and inner layers and fitler material on the middle layer.

7. ESSENTIAL HEALTH AND SAFETY REQUIREMENTS

- A visual inspection was made according to EN 149:2001 +A1:2009 for ergonomics.
- Protection levels and degrees are defined by the manufacturer.
- Suitable construction materials were determined by visual inspection according to EN 149:2001 +A1:2009.

8. ANALYSIS EVALUATION AND MARKING:

EN 149:2001 +A1:2009

1	TESTS	PARAMETER	PERFORMANCE LEVELS		ICE	RESULTS	PERFORMAN CE LEVELS	EVALUATIO N
			FFP1	FFP 2	FFP3			
	Part 7.3	Shall also the marking and the information supplied by the manufacturer				Appropriate	-	PASS

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CHECK AT RANDOM INTERVALS

Visual				
inspection				
Banned Azo	< 30 mg/kg	< 5 mg/kg	-	PASS
Dyes				
Part 7.4	Particle filtering half mask shall be offered for	Appropriate	-	PASS
Packaging	sale packaged in such a way that they are			
	protected against mechanical damage and			
	contamination before use.			
Part 7.5	When conditioned in accordance 8.3.1 &	Appropriate	-	PASS
Material	8.3.2 the particle filter half mask shall not			
	collapse.			
Part 7.6	After cleaning and disinfecting the re-usable	Not applicable	-	Not
Cleaning and	particle filtering half mask shall satisfy the			applicable
disinfecting	penetration requirement of the relevant			
	class.			
Part 7.7	No negative comments should be made by	Appropriate	-	PASS
Practical	the test subject regarding any of the criteria			
performance	evaluated.			
Part 7.8	Parts of the device likely to come into contact	Appropriate	-	PASS
Finish of parts	with the wearer shall have no sharp edge or			
	burrs.			

TESTS	PARAMETER	PERFORMANCE LEVELS		RESULTS	PERFORMANCE LEVELS	EVALUATION	
		FFP1	FFP2	FFP3			
Part 7.9.1	At least 46 out of the	≤25	≤11	≤5	-	-	-
Total inward	50 individual						
leakage	exercise result						
	At least 8 out of the	≤22	≤8	≤2	-	-	-
	10 individual wearer						
	arithmetic means						

Subject facial dimensions

Subject	Face Length (mm)	Face Width (mm)	Face Depth (mm)	Mouth Width (mm)
				+ \
1	133	132	132	65
2	125	144	116	67
3	126	135	124	75
4	123	133	134	74
5	117	135	122	73
6	122	142	133	66
7	113	132	114	75
8	135	123	123	65
9	122	135	133	74
10	135	142	125	83

TESTS	PARAMETER	PERFORMANCE LEVELS		RESULTS	PERFORMANCE LEVELS	EVALUATION	
		FFP1	FFP2	FFP3			
Part 7.9.2	Sodium chloride, 95	% 20	% 6	% 1	See the	FFP3	PASS
Penetration	L/min			1	table below		
of filter	%, max	X	\ \ \ \	1/1/1			
material	Paraffin oil, 95 L/min	% 20	% 6	% 1	See the	FFP3	PASS
	%, max				table below		



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Penetration of filter material	Sodium Chloride (%)	Paraffin Oil (%)
As received	0,1	0,2
As received	0,3	0,4
As received	0,2	0,2
After the simulated wearing treatment	0,3	0,4
After the simulated wearing treatment	0,2	0,4
After the simulated wearing treatment	0,3	0,3
Mechanical strength and temperature conditioning	0,4	0,7
Mechanical strength and temperature conditioning	0,5	0,6
Mechanical strength and temperature conditioning	0,4	0,6

TESTS	PARAMETER PERFORMANCE		RESULTS	PERFORMANCE	EVALUATION		
	LEVELS			LEVELS			
		FFP1	FFP2	FFP3			
Part 7.10	Materials shall not b	e know	n to be	likely to	Appropriate	-	PASS
Compatibility	cause irritation or an	y other a	adverse	effect to			
with skin	health						
Part 7.11	Mask shall not burn o	or not to	continue	e to burn	-	-	-
Flammibility	for more than 5 s						
Part 7.12	Shall not exceed an	average	of % 1		-	-	-
Carbondioxide							
content of the							
inhalation air							
Part 7.13	It can be donned and	d remove	ed easily	У	Appropriate	-	PASS
Head harness							
Part 7.14	The field of vision sha	all accep	table in	practical	Appropriate	-	PASS
Field of vision	performance test.						
Part 7.15	It shall withstand axi	ally a te	nsile for	ce of 10	Not	-	Not applicable
Exhalation	N apply for 10 s.				applicable		
valve(s)	If fitted, shall contin						
	after a continuous		on flow	ot 300			
	L/min over a period of	ภ 3U S.					

TESTS	PARAMETER	PERFORMANCE LEVELS		RESULTS	PERFORMANCE LEVELS	EVALUATION	
		FFP1	FFP2	FFP3			
Part 7.16	Inhalation 30L/min	0,6	0,7	1,0	See the table	FFP3	PASS
Breathing		mbar	mbar	mbar	below		
Resistance	Inhalation 95L/min	2,1	2,4	3,0	See the table	FFP3	PASS
		mbar	mbar	mbar	below		
	Exhalation	3,0	3,0	3,0	See the table	FFP3	PASS
	160L/min	mbar	mbar	mbar	below		

Breathing Resistance (mbar)	Inhalation 30L/min	Inhalation 95L/min
As received	0,4	1,5
As received	0,4	1,5
As received	0,4	1,5
After temperature conditioning	0,3	1,5
After temperature conditioning	0,3	1,4
After temperature conditioning	0,3	1,4
After the simulated wearing treatment	0,3	1,4
After the simulated wearing treatment	0,3	1,4
After the simulated wearing treatment	0,4	1,4
After the flow conditioning	V - /	1 - / /
After the flow conditioning		-
After the flow conditioning	1 -	- \



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Breathing Resistance 160L/min (mbar)	Facing directly ahead	Facing vertically upwards	Facing vertically downwards	Lying on the left side	Lying on the right side
As received	2,4	2,4	2,4	2,3	2,4
As received	2,4	2,4	2,4	2,4	2,4
As received	2,4	2,3	2,4	2,4	2,3
After temperature conditioning	2,4	2,3	2,3	2,4	2,3
After temperature conditioning	2,4	2,3	2,4	2,3	2,3
After temperature conditioning	2,4	2,3	2,3	2,4	2,3
After the simulated wearing treatment	2,4	2,3	2,4	2,4	2,4
After the simulated wearing treatment	2,4	2,3	2,3	2,4	2,3
After the simulated wearing treatment	2,3	2,3	2,4	2,3	2,4
After the flow conditioning	-	-	-	-	-
After the flow conditioning	-	-	-	-	-
After the flow conditioning	-	-	-	-	-

TESTS	PARAMETER	PERFORMANCE		RESULTS	PERFORMANCE	EVALUATION	
			LEVELS			LEVELS	
		FFP	FFP	FFP3			
		1	2				
Part 7.17	After clogging the	4	5	7	Not applicable	-	Not applicable
Clogging	inhalation	mbar	mbar	mbar			
	resistances shall						
	not exceed.						
	(valved)						
	The exhalation resist	ance sh	all not	exceed	Not applicable	-	Not applicable
	3 mbar at 160 L/	min co	ntinuou	s flow.			
	(valved)						
	After clogging the	3	4	5	Not applicable	-	Not applicable
	inhalation and	mbar	mbar	mbar			
	exhalation						
	resistances shall						
	not exceed.						
	(valveless)						
Part 7.18	All demountable par	rts (if f	itted) sl	hall be	Not applicable	-	Not applicable
Demountable	readily connected	and s	secured	were			
part	possible by hand.						
Part 9	The packaging inform				Appropriate	-	PASS
Marking	and durably marke						
	commercially availab						
	through it if the packa	aging is	transpa	rent.			

9. ATTACHMENTS

Test Reports (M-2024-01327, M-2024-01358)

CONTROLLER

SIGNATURE

DATE



Report Nu.: M-2024-01327 Date: 2024-10-09 16:57:45 Page: 1 / 4 Rev:

Purpose of Analysis : Special request

Sample Send Org. : SJT-Investment Group Oy / JedX Medcare

Address : Köysikuja 1, 01640 Vntaa, Finland

Sample Acceptance Date : 2024-10-02 17:55:32

Analysis Date : 2024-10-02 00:35:17

Sample Quantity : 120 Pieces

Sample Description : JedX 1625 W HL

Other informations :

Tests	Method	Expected performance level	Evaluation
Penetration Of Filter Material	EN 149+A1 Part 8.11, EN 13274-7		PASS (FFP3)
Breathing Resistance	EN 149+A1 Part 8.9		PASS (FFP3)

Penetration Of Filter Material

Device:Filter Test System

Measurement uncertainty:±0,080

Tests	Analysis result	Limit Value	Method	Evaluation	Physical Condition
Penetration Of Filter Material	Check the table.	FFP1≤20 FFP2≤6 FFP3≤1	EN 149+A1 Part 8.11, EN 13274-7	PASS (FFP3)	-

	Sodium Chloride (%)	Paraffin Oil (%)
As received 1	0,1	0,2
As received 2	0,3	0,4
As received 3	0,2	0,2
After the simulated wearing treatment 1	0,3	0,4
After the simulated wearing treatment 2	0,2	0,4
After the simulated wearing treatment 3	0,3	0,3
Mechanical strength and temperature conditioning (120 mg) 1	0,4	0,7
Mechanical strength and temperature conditioning (120 mg) 2	0,5	0,6
Mechanical strength and temperature conditioning (120 mg) 3	0,4	0,6



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1 '		,	

Breathing Resistance

Device:Breathing Resistance Tester

Measurement uncertainty: Inhalation 30L/min:±0,160,Inhalation30 L/min:±0,026 Exhalation 160 L/min:0,046

Tests	Analysis result	Limit Value	Method	Evaluation	Physical Condition
Breathing Resistance	Check the table.	See the limits table.	EN 149+A1 Part 8.9	PASS (FFP3)	-

Classification	30 L/min max basınç (mbar)	95 L/min max basınç (mbar)	160 L/min max basınç (mbar)
FFP1	0,6	2,1	3,0
FFP2	0,7	2,4	3,0
FFP3	1,0	3,0	3,0

Inhalation	30 L/min	95 L/min	
As received 1	0,4	1,5	
As received 2	0,4	1,5	
As received 3	0,4	1,5	
After temperature conditioning 1	0,3	1,5	
After temperature conditioning 2	0,3	1,4	
After temperature conditioning 3	0,3	1,4	
After the simulated wearing treatment 1	0,3	1,4	
After the simulated wearing treatment 2	0,3	1,4	
After the simulated wearing treatment 3	0,4	1,4	
After the flow conditioning 1	-	-	
After the flow conditioning 2	-	-	
After the flow conditioning 3			

Exhalation 160L/min	Facing directly ahead	Facing vertically upwards	Facing vertically downwards	Lying on the left side	Lying on the right side
As received 1	2,4	2,4	2,4	2,3	2,4
As received 2	2,4	2,4	2,4	2,4	2,4
As received 3	2,4	2,3	2,4	2,4	2,3
After temperature conditioning 1	2,4	2,3	2,3	2,4	2,3



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After temperature conditioning 2	2,4		2,3	2,4	2,3	2,3
After temperature conditioning 3	2,4		2,3	2,3	2,4	2,3
After the simulated wearing treatment 1	2,4		2,3	2,4	2,4	2,4
After the simulated wearing treatment 2	2,4		2,3	2,3	2,4	2,3
After the simulated wearing treatment 3	2,3		2,3	2,4	2,3	2,4
After the flow conditioning 1	-		-	-	-	-
After the flow conditioning 2	-		-	-	-	-
After the flow conditioning 3						



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- 4. Results are valid for the sample received.
- 5. The decision rule is the rule that determines how measurement uncertainty is taken into account when specifying compliance with an established specification. The customer may choose to apply and/or not apply the decision rule (except in cases where legislation/standards are mandatory). If the customer prefers to apply the decision rule; According to the TLM-052 Decision Rule Application instruction published on the www.mnalab.com website, the decision rule selected in agreement is applied and reported by stating the relevant analysis and decision rule method in the "Note" section. If the customer leaves the decision rule application to the laboratory's evaluation, MNA LABORATORIES applies the simple decision rule.
- 6. Limit Values are determined by taking from analysis methods.
- 7. The laboratory is not responsible if the information provided by the CUSTOMER affects the validity of the results.
- 8. Test and / or measurement results, expanded measurement uncertainties (if any) and test methods are given in the following pages, which are the supplementary part of this certificate.

The witness sample not recieved.

Nesligül Arkan

Sample Acceptance and Reporting Officer

2024-10-09 16:57:33

Erhan Üstünel

Laboratory Responsible

2024-10-09 16:57:02

ΥΟΙ ΚΑΝ ΔΚΙΝ

Laboratory Manager

2024-10-09 16:57:13



ANALYSIS REPORT



AB-1183-T

M-2024-01358

10-24

Report Nu.: M-2024-01358 Date: 2024-10-09 08:57:57 Page: 1 / 2 Rev:

Purpose of Analysis : Special request

Sample Send Org. : SJT-Investment Group Oy / JedX Medcare

Address : Köysikuja 1, 01640 Vntaa, Finland

Sample Acceptance Date : 2024-10-07 08:19:50

Analysis Date : 2024-10-07 08:34:18

Sample Quantity : 4 Pieces

Sample Description : Jedx black and green

Other informations :

Tests	Method	Expected performance level	Evaluation
Banned Azo Dyes	Banned Azo Dyes EN ISO 14362-1 / EN ISO 17234-1		PASS

Banned Azo Dyes *

Device:GC-MS

Measurement uncertainty: Textile:±0,350 Leather:±0,390

2-naphthylamine/91-59-8, o-aminoazotoluene, 4-amino-2',3-dimethylazobenzene, 4-o-tolylazo-o-toluidine/97-56-3, 5-nitro-o-toluidine/99-55-8, 4-chloroaniline/106-47-8, 4-methoxy-m-phenylenediamine/615-05-4, 4,4'-methylenedianiline, 4,4'-diaminodiphenylmethane/101-77-9, 3,3'-dichlorobenzidine, 3,3'-dichlorobiphenyl-4,4'-ylenediamine/91-94-1, 3,3'-dimethoxybenzidine, o-dianisidine/119-90-4, 3,3'-dimethylbenzidine, 4,4'-bi-o-toluidine/119-93-7, 4,4'-methylenedia-o-toluidine/838-88-0, 6-methoxy-m-toluidine p-cresidine/120-71-8, 4,4'-methylene-bis-(2-chloro-aniline), 2,2'-dichloro-4,4'-methylene-dianiline/101-14-4, 4,4'-oxydianiline/101-80-4, 4,4'-thiodianiline/139-65-1, o-toluidine, 2-aminotoluene/95-53-4, 4-methyl-m-phenylenediamine/95-80-7, 2,4,5-trimethylaniline/137-17-7, o-anisidine, 2-methoxyaniline/90-04-0, 4-amino azobenzene/60-09-3

Compounds/Cas No: biphenyl-4-ylamine, 4-aminobiphenyl xenylamine/92-67-1, benzidine/92-87-5, 4-chloro-o-toluidine/95-69-2,

Tests	Analysis result	Limit Value	Method	Evaluation	Physical Condition
Banned Azo Dyes	Check the table.	≤30 mg/kg	EN ISO 14362-1 / EN ISO 17234-1	PASS	-

Part of Sample/ Numune kısımları	Results/ Sonuçlar(mg/kg)	
Black Fabric+Black elastic earloop+Green elastic earloop	<5	





AB-1183-T

M-2024-01358

10-24

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Operating as a test laboratory, MNA Laboratories is accredited by TÜRKAK according to AB-1183-T and TS_EN_ISO/IEC_17025:2017 standards has been done. A multilateral agreement with the European Accreditation Association (EA) on the recognition of the Turkish Accreditation Agency (TÜRKAK) test reports and It has signed a mutual recognition agreement with the International Laboratory Accreditation Association (ILAC).

*The analysis is within the scope of accreditation.

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- 7. The laboratory is not responsible if the information provided by the CUSTOMER affects the validity of the results.
- 8. Test and / or measurement results, expanded measurement uncertainties (if any) and test methods are given in the following pages, which are the supplementary part of this certificate.

The witness sample was not received.

Nesligül Arkan

Sample Acceptance and Reporting Officer

2024-10-09 08:57:12

Erhan Üstünel Laboratory Responsible

2024-10-09 08:56:16

VOLKAN AKIN

Laboratory Manager

2024-10-09 08:56:26

Suns